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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/539,505	01/09/2006	Joerg Rosenberg	M/43212-US-1	4705
26474 NOVAK DRI	7590 06/15/201 ICE DELUCA + QUIG	EXAM	EXAMINER	
300 NEW JER	SEY AVENUE NW	KATAKAM, SUDHAKAR		
FIFTH FLOOI WASHINGTO		ART UNIT	PAPER NUMBER	
			1621	
			MAIL DATE	DELIVERY MODE
			06/15/2010	PAPER

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Office Action Summary

A P				
Application No.	Applicant(s)			
10/539,505	ROSENBERG ET AL.			
Examiner	Art Unit			
SUDHAKAR KATAKAM	1621			

-- The MAILING DATE of this communication appears on the cover sheet with the correspondence address --Period for Reply

A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) OR THIRTY (30) DAYS,

- WHICHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION.
- Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed
- after SIX (6) MONTHS from the mailing date of this communication. - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication.
- Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133).

 Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any

earned pater	t term adjustment.	See 37 CFR	1.704(b).	

Status					
1)🛛	Responsive to communication(s) filed on <u>31 March 2010</u> .				
2a)□	This action is FINAL . 2b)⊠ This action is	non-final.			
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is				
	closed in accordance with the practice under Ex parte 0	Quayle, 1935 C.D. 11, 453 O.G. 213.			
Disposit	ion of Claims				
4)🖂	Claim(s) 23-31 and 33-38 is/are pending in the applicat	on.			
~	4a) Of the above claim(s) is/are withdrawn from consideration.				
5)	Claim(s) is/are allowed.				
6)🖂	Claim(s) 23-31 and 33-38 is/are rejected.				
7)	Claim(s) is/are objected to.				
8)□	Claim(s) are subject to restriction and/or election	requirement.			
Applicat	ion Papers				
91□	The specification is objected to by the Examiner.				
	The drawing(s) filed on is/are: a) accepted or	b) Objected to by the Examiner.			
,	Applicant may not request that any objection to the drawing(s				
	Replacement drawing sheet(s) including the correction is requ				
11)	The oath or declaration is objected to by the Examiner.	• • • • • • • • • • • • • • • • • • • •			
Priority (under 35 U.S.C. § 119				
12)🖾	Acknowledgment is made of a claim for foreign priority u	nder 35 U.S.C. § 119(a)-(d) or (f).			
a)⊠ All b)□ Some * c)□ None of:					
	 Certified copies of the priority documents have been received. 				
	2. Certified copies of the priority documents have been received in Application No				
	3. Copies of the certified copies of the priority documents have been received in this National Stage				
application from the International Bureau (PCT Rule 17.2(a)).					
* (See the attached detailed Office action for a list of the ce	rtified copies not received.			
Attachmen	t(s)				
	ce of References Cited (PTO-892)	4) Interview Summary (PTO-413)			
	e of Draftsperson's Patent Drawing Review (PTO-948)	Paper No(s)/Mail Date 5) Notice of Informal Patent Application.			
	mation Disclosure Statement(s) (FTC/SB/08) or No(s)/Mail Date	6) Other:			
S. Patent and T	redemark Office (ev. 08-06) Office Action Sums	part of Paper No./Mail Date 20100612			
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DETAILED ACTION

Status of the application

 Receipt of Applicant's request for continued examination filed on 31 March 2010 is acknowledged.

A request for continued examination under 37 CFR 1.114, including the fee set forth in 37 CFR 1.17(e), was filed in this application after final rejection. Since this application is eligible for continued examination under 37 CFR 1.114, and the fee set forth in 37 CFR 1.17(e) has been timely paid, the finality of the previous Office action has been withdrawn pursuant to 37 CFR 1.114. Applicant's submission filed 31 March 2010 has been entered.

2. Claims 23-31 and 33-37 are examined on the merits in this office action.

Claim Rejections - 35 USC § 103

 The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this little, if the differences between the subject matter sought to be patented and the prior at are such that the subject matter sought to the patented and the prior at are such that the subject matter possible of the subject matter possible sought to the patents. Patentially shall not be negatived by the manner in which the invention was made.

- 4. The factual inquiries set forth in *Graham* v. *John Deere Co.*, 383 U.S. 1, 148 USPQ 459 (1966), that are applied for establishing a background for determining obviousness under 35 U.S.C. 103(a) are summarized as follows:
 - Determining the scope and contents of the prior art.
 - Ascertaining the differences between the prior art and the claims at issue.
 - 3. Resolving the level of ordinary skill in the pertinent art.
 - Considering objective evidence present in the application indicating obviousness or nonobviousness

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 Claims 23-31 and 33-37 are rejected under 35 U.S.C. 103(a) as being unpatentable over by Boyer et al (US 4,800,079) in view of Kothrade et al (US 6,284,803).

Boyer teaches a medicine based on fenofibrate, and a method of preparing it.

Boyer defined the term "fenofibrate and its derivatives" to designate a compound having the formula I, is represented by the following formula:

$$R_1$$
-co- $\begin{pmatrix} R_2 \\ CH_3 \\ CH_3 \end{pmatrix}$

The above formula reads instant claim 1 when R_1 is phenyl group, R_2 and R_3 are hydrogen atoms, and Y is a -OH group [col. 1, lines 10-31]. **Boyer also** teaches various binders, selected from the group comprising methacrylic polymers, polyvinylpyrolidone, mixtures thereof; cellulose derivatives and polyethylene glycols (see claim 2).

Boyer et al is deficient in sense that the dependent limitations in the claims 24-31 and 33-37 are not explicitly stated in the composition. However, Kothrade et al cure this deficiency.

Kothrade et al teach a pharmaceutical formulation [col. 14, line 45] in dosage form [col. 1, line 4] comprising fenofibrate as the active ingredient [col. 7, line 39], in the form of a molecular dispersion [col. 10, line 48], and a polymeric

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binder composed of methy/methacrylate, acrylic acid, cellulose acetate phthalate and hydroxypropylmethylcellulose phthalate [col. 5, lines 11-13, 20-21] and other conventionally acceptable excipients [col. 1, lines 4-7], which include flow regulators and silicates/silica gel [col. 6, lines 1 and 12]. The formulation is further obtainable by melt extrusion [col. 2, line 8; col. 5, line 35]. The formulation has a ratio of free carboxyl groups to esterified carboxyl groups around 1:1, based on the weight percentage of methyl methacrylate to acrylic acid [col. 2, lines 56-61] and the use of Eudragit types, which Applicant uses to exemplify this ratio preference [col. 5, line 12; col. 10, line 391 [see also specification page 7, lines 3-10]. The formulation comprises 0.1 to 95%, preferably from 20 to 80%, in particular 30 to 70% by weight of the active substance [col. 6, lines 61-63], with ranges of 15-83% for the binder [col. 2, lines 19-45], in which the enteric binder (Eudragit types) is in the preferable range of up to 75% by weight of the binder component [col. 4, lines 65-67; col. 5, line 1 and 12] and with the range of up to 100%, in particular 0.02-50% of pharmaceutically/physiologically acceptable additives [col. 5, lines 66-67; col. 6, lines 7-81. The preceding percentages would include a formulation in which the content of active substance component relative to binder is from 20 to 30% by weight.

Kothrade et al further teaches that all three components of the formulation: fenofibrate, binder component and other excipients/additives, can be combined [col. 1. lines 4-7; col. 7. lines 10-12 and 39].

With regard to claim 37, which describes a method for oral administration, since the dosage is in tablet form [col. 10, line 67], the expected mode of administration is oral

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administration. Additionally, applicant states that fenofibrate is usually administered orally [specification page 1, line 15].

With regard to claim 25 and 26, which describes the binder as an enteric binder/enteric polymer, because the art describes the polymeric binder with the same components as applicant's, which include methyl methacrylate, acrylic acid, cellulose, acetate phthalate and hydroxypropylmethylcellulose phthalate [col. 5, lines 11-13, 20-21], therefore, the enteric property is inherent to the binder/polymer composition.

The claim would have been obvious because the design incentives or market forces provided a reason to make an adaptation, and the invention resulted from application of the prior knowledge in a predictable manner.

All the claimed elements were known in the prior art and one skilled person in the art could have combined the elements as claimed by known methods with no change in their respective functions, and the combination would have yielded predictable results to have yielded predictable results to one of ordinary skill in the art at the time of the invention.

Therefore, it would be prima facie obvious to one of ordinary skill in the art at the time of the invention, to combine the components of **Kothrade et al** for the formulation of **Boyer et al** to arrive at a fenofibric acid composition for pharmaceutical oral administration. The expected result would be an effective lipid-regulating tablet in dosage form.

Response to Arguments

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Applicant's arguments filed on 31 March 2010 have been fully considered but they are not persuasive.

The examiner acknowledges applicants argument that "it has been observed that fenofibrate has poor solubility in aqueous liquids, thereby giving rise to non-uniform absorption in the digestive tube, and in accordance with the present invention a galenical preparation has been devised which considerably improve absorption by the digestive tube.

Boyer, defined the meaning of "fenofibrate and its derivatives". The **Boyer's** formula (I) becomes fenofibric acid, when R_1 is phenyl group, R_2 and R_3 are hydrogen atoms, and Y is a -OH group [col. 1, lines 10-31]. However, Boyer fails to exemplify a formulation with fenofibric acid in their disclosure.

Fenofibric acid is known in the art. The acid or salt form of a compound is preferable over its ester form in the formulations. In this case, fenofibric acid is expected to have high solubility over its ester form and will have better absorption properties over the fenofibrate. The claim would have been obvious because the design incentives or market forces provided a reason to make an adaptation, and the invention resulted from application of the prior knowledge in a predictable manner.

Therefore, it would be prima facie obvious to one of ordinary skill in the art at the time of the invention, to combine the above cited references and arrive at a fenofibric acid composition for pharmaceutical oral administration with a reasonable expectation of success. The expected result would be an effective lipid-regulating tablet in dosage form.

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Conclusion

7. No claim is allowed.

 Any inquiry concerning this communication or earlier communications from the examiner should be directed to Sudhakar Katakam whose telephone number is 571-272-9929. The examiner can normally be reached on M-F 8:30 AM - 5:00 PM.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Daniel Sullivan can be reached on 571-272-0779. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/Sudhakar Katakam/

Examiner, Art Unit 1621